Materials and technologies for fabrication of three-dimensional microstructures with sub-100 nm feature sizes by two-photon polymerization

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Materials and technologies for fabrication of three-dimensional microstructures with sub-100 nm feature sizes by two-photon polymerization

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The fabrication of sub-100 nm feature sizes in large-scale three-dimensional (3D) geometries by two-photon polymerization requires a precise control of the polymeric reactions as well as of the intensity distribution of the ultrashort laser pulses. The authors, therefore, investigate the complex interplay of photore sist, processing parameters, and focusing optics. New types of inorganic–organic hybrid polymers are synthesized and characterized with respect to achievable structure sizes and their degree of crosslinking. For maintaining diffraction-limited focal conditions within the 3D processing region, a special hybrid optics is developed, where spatial and chromatic aberrations are compensated by a diffractive optical element. Feature sizes below 100 nm are demonstrated. © 2012 Laser Institute of America.

Key words: two-photon polymerization, hybrid polymer, ORMOCER, refractive-diffractive hybrid optics, aberration correction

I. INTRODUCTION

In recent years, two-photon polymerization (2PP) has proven to be technology capable of fabricating arbitrary three-dimensional (3D) microstructures with feature sizes smaller than the diffraction limit of the applied laser wavelength. Developments of new materials and illumination concepts like the STED lithography have pushed the lateral dimensions of realizable structures below 100 nm. With the variability of the technology, several applications have emerged, e.g., in the field of micro-optics, where the fabrication of photonic crystals or micro-optical elements has been demonstrated. Furthermore, biomedical devices like scaffolds for cell growth or small prostheses could be realized.

Despite the ongoing progress in the field of 2PP, it remains challenging to preserve sub-100 nm feature sizes for complex three-dimensional microstructures with several 10 µm in height. This is mainly due to structuring depth dependent aberrations that occur when femtosecond laser pulses are focused with conventional microscope objectives into a nonindex-matched polymer. These aberrations lead to a decrease of the peak intensity in the focal plane and consequently to an inhomogeneous structuring process for different writing depths.

Besides the focusing optics and the applied process parameters, the molecular structure of the employed photopolymer plays an important role in structure formation as well as the achievable spatial resolution and surface roughness of the cured structures.

Therefore, we investigate the impact of material, process parameters, and focusing optics. Specially, adapted new types of the inorganic–organic hybrid polymer ORMOCER® were synthesized and characterized with respect to achievable feature sizes and degree of crosslinking (Sec. II). This gives a detailed insight into the impact of crosslinking chemical groups, photoinitiators, and laser parameters (pulse energy, exposure time) on structure formation. Furthermore, a refractive-diffractive hybrid optics for aberration-free focusing of the laser pulses into the polymer was developed and improvements for the volume structuring are presented (Sec. III).

II. MATERIAL SYNTHESIS AND PATTERNING

ORMOCER®'s have become a material class of great interest for 2PP, since they combine favorable properties of glasslike materials, such as a high mechanical, thermal, and chemical stability with the ability to be photochemically patterned, which is well known from purely organic photore sist. Furthermore, they exhibit excellent optical properties and biocompatibility, thus being a perfect material for micro-optical and biomedical applications. For synthesis, alcoxysilane-precursors undergo hydrolysis and polycondensation reactions resulting in an organically modified inorganic-oxidic network (Si–O)n. This liquid resin can then be crosslinked photochemically and/or thermally resulting in solidified structures. As the reaction product
strongly depends on the choice of precursors and synthesis conditions, various properties of ORMOCER®s like stability, refractive index, and absorption behavior, as well as their molecular structure and polymerizable moiety can be tailored. Thus, there are several types of ORMOCER®s, each one adapted to one or more specific applications.

For efficient organic crosslinking of ORMOCER®s using 2PP suitable photoinitiators had to be identified, which show a high two-photon absorption cross-section, combined with a high quantum efficiency. Z-Scan experiments carried out on several commercial as well as specifically designed initiators revealed these conditions being met by some substances like the Ciba Irgacure® product line.27,28 To investigate the resulting structure sizes for different material formulations and laser parameters, ascending scan experiments29 were conducted on the methacrylic and acrylic ORMOCER®s OC-I and OC-V, respectively. In these studies, it turned out that in both ORMOCER® systems the achievable structure sizes did not reach the 100 nm threshold.27

One drawback of exploiting the material’s threshold behavior in order to improve the spatial resolution is that the degree of polymer conversion, i.e., the degree of organic crosslinking, decreases dramatically when using lower photon fluxes. This affects the hardness of the solidified volume and, thus, the stability of 3D structures. A similar effect occurs when aberrations lead to a decrease of the peak intensity inside the volume of the polymer. The stability of voxels fabricated in the volume would diminish compared to voxels on the substrate. In order to quantify this loss of mechanical stability, we performed μ-Raman investigations on 2PP fabricated cube structures. These cubes were written in OC-I with different photon fluxes and hatching distances. We used a frequency-doubled Yb:YAG oscillator (t-Pulse 200, Amplitude Systemes) with a fundamental laser wavelength of 1030 nm, a pulse duration of 400 fs, and a repetition rate of 10 MHz in our experimental setup (described in detail in the literature).4 To deduce the degree of polymer conversion, K, from the μ-Raman spectra, one needs a reference peak, which does not change upon illumination, and a peak of the cross-linking groups. In case of OC-I, the reference peak at 1570 cm\(^{-1}\) stems from ring vibrations of diphenylsilane-diol (DPD), whereas the C=C peak at 1640 cm\(^{-1}\) is an indicator for the organic crosslinking of the material. With the formula

\[ K = 100 \times \left[ 1 - \frac{A_{1570}}{A_{1640}} \frac{A_{1640}}{A_{1570}} \right], \]

K can be calculated,24 where A denotes the area of the respective peaks of the polymerized structure and A’ is the liquid ORMOCER® resin. Due to the overlapping of several peaks from initiator and ORMOCER® in the spectral region of interest, a deconvolution of peaks has been applied using the commercial software Omnic from Thermo Scientific. Two exemplary spectra of nonpolymerized and polymerized materials are shown in Fig. 1. As demanded, the peak at 1570 cm\(^{-1}\) is not affected after 2PP, whereas the peak at 1640 cm\(^{-1}\) diminishes. The results of our μ-Raman experiments are displayed in Fig. 2. As expected, the polymer conversion increases with increasing laser power until it reaches saturation at comparatively high laser powers. A further increase of power then does no longer affect the degree of crosslinking. The saturation value of approximately 80% is slightly higher compared to results found in thin layers of OC-I solidified using mask-aligner i-line illumination. Cubes with a K smaller than 35% were mechanically unstable and did not survive the solvent wash after illumination. This has to be kept in mind in the quest for 3D sub-100 nm 2PP lithography.

One aspect which is not yet entirely understood is the dependency of K on the hatching distance. The primary crosslinking of a single rod created during hatching of a cube should only be depending on the applied photon flux. But since we measured entirely polymerized cubes (rather than crystal structures with voids), single rods could not be measured. Actually, we average over overlapping rods, since their width is expected to be at least equal to or probably larger than the hatching distance. Furthermore, it has to be mentioned that even without the overlapping of rods, the measurement spot of the instrument always integrates over some rods, since its spatial resolution is, in contrast to 2PP, diffraction-limited. Therefore, the smaller distance of adjacent rods leads to measurement of higher conversion because of these two effects. This also agrees with the intuitive guess that structures with smaller hatching distances must be crosslinked stronger, because their total exposure time to laser pulses was longer, corresponding to higher exposure dose. Another issue might be the lifetime of radicals responsible for the crosslinking. Radicals created while writing a first rod might still influence the creation of the adjacent rod, if
their lifetime is long enough and the adjacent rod is not too far away. This is subject of further investigations. In order to obtain a better understanding of crosslinking during the polymerization process, simulations of the molecular structure of the OC-I monomers and polymerization products are currently performed.30

Based on the findings from these preliminary experiments, a new ORMOCER® system, OC-II, was synthesized and patterned. In analogy to OC-I, it was synthesized from DPD and p-styryl-trimethoxysilane, i.e., the main difference is the polymerizable group, which is styryl in OC-II. Figure 3(a) depicts results of ascending scan experiments for average laser powers from 200 to 500 μW. Structure sizes below 100 nm can be achieved easily. In order to reveal the underlying scaling mechanism, Fig. 3(b) shows voxels of constant diameter plotted versus applied laser power and exposure time. We assumed an exposure dose \( D \propto P^N t \) with \( P \) being the laser power and \( t \) the exposure time. A constant exposure dose \( D \) results in constant feature sizes. The dashed line corresponds to linear absorption \( (N = 1) \) and the solid line to a two-photon absorption process \( (N = 2) \). The good agreement of the slope of the solid line with the measured data indicates that two-photon absorption induced polymerization is the dominant process in structure formation. However, a possible contribution of thermal effects on the polymerization process caused by the high repetition rate of the laser system31 has to be investigated in further experiments.

In Fig. 4, scanning electron microscope (SEM) images of some example structures are shown revealing 91 nm voxel sizes and line widths in the order of 100 nm. It is likely that the reactivity of the styryl group in comparison to the methacryl group in OC-I leads to the improvement in feature size, although this has to be investigated in further detail as well.

III. OPTICAL DESIGN

In order to fabricate stable microstructures with a high degree of crosslinking and sub-100 nm feature sizes, aberration-free focusing of ultrashort laser pulses into the polymer for the complete structure height is needed. Usually, oil immersion microscope objectives with high numerical apertures (NAs) up to 1.45 are applied in setups for 2PP.12,15 The laser pulses are focused through index-matching oil and a coverslip into the polymer located on the bottom of the coverslip. These conventional microscope objectives suffer from two main drawbacks when applied for 2PP. Diffraction-limited focusing of the laser pulses requires that the design conditions of the objectives are met, which are thickness and refractive index of the coverslip and usage of the proper immersion oil. Already a small change of the refractive index between coverslip and polymer leads to strong spherical aberration, whose amount mainly depend on the difference in the refractive indices, the focusing depth, and the NA of the objective.32,33 Second, microscope objectives are complex optical systems consisting of several lenses causing a high group velocity dispersion (GVD), which leads to a temporal broadening of the laser pulses. In order to maintain the initial pulse duration, external compensation of the GVD is needed.

To overcome these drawbacks, we designed and manufactured a high numerical aperture hybrid optics specifically for the structuring of the ORMOCER® OC-I via 2PP. However, the optics can be easily adapted to other polymers (e.g., OC-II) as well.

The optics consists of merely three optical elements in order to keep the internal dispersion to a minimum: a diffractive optical element (DOE) for achromatization and the correction of spherical aberrations,34 an asphere with NA = 0.6 (Edmund Optics 47 093), and a half ball lens working as a solid immersion lens (SIL) to increase the NA. The design of the hybrid optics allows for diffraction-limited focusing of the ultrashort pulses into the polymer with a numerical aperture NA = 1.33 over a working distance range of 550 μm. The optics is designed for focusing ultrashort pulses with a central wavelength of 515 nm and a spectral width of 3 nm.
A SIL offers the possibility to significantly increase the NA of the asphere without introducing additional aberrations. There are two possible configurations for the SIL. In the case of a hemispherical SIL, a hemisphere made of glass with refractive index $n$ is placed with its center of curvature in the focus of the objective. When the rays enter perpendicular to the surface normal, no refraction occurs and the NA of the objective is increased by a factor $n$. Our design comprises an aplanatic solid immersion lens (ASIL). The ASIL consists of a hyper-hemisphere with radius $R$ and refraction index $n$. For a center thickness of $d = R(1 + \frac{1}{n})$, all geometrical aberrations from the ASIL are equal to zero. The NA of the objective is increased by a factor $n^2$, due to the additional refraction at the surface of the ASIL. For three-dimensional structuring of the polymer, the ASIL has to be integrated into the setup in such a way that aberration-free focusing for variable focusing depths is possible. As shown in Fig. 5, the ASIL is composed of a half ball lens, immersion liquid, substrate and polymer. By choosing the polymer without the photoinitiator as immersion liquid, the total optical path inside the polymer remains constant for different focusing depths over the complete working distance. However, this setup introduces small spherical aberrations, since the ASIL no longer consists of one material with homogeneous refractive index. They can be minimized by choosing a glass for the half ball lens with a refractive index very close to that of the polymer. In our case, the half ball lens is made of N-SK11 with $n_d = 1.564$, almost exactly coinciding with $n_d = 1.563$ of the polymer.

For characterization, we compare our optics with an oil immersion microscope objective (Plan-Apochromat 100x, NA = 1.4, immersion oil Immersol $n_e = 1.518$, Zeiss) in terms of the structure sizes directly on the substrate surface as well as for different writing depths inside the polymer. In our experiments, a frequency-doubled Yb:YAG oscillator (t-Pulse 500, Amplitude Systemes) with a fundamental laser wavelength of 1030 nm, a pulse duration of 500 fs, and a repetition rate of 10 MHz is used. An acousto-optic modulator (AOM) is applied as shutter; the laser power is adjusted with a polarization beam splitter in combination with a half-wave plate.

For structuring directly on the substrate surface, the microscope objective as well as the hybrid optics are designed for diffraction-limited focusing. We applied point-by-point illumination of the polymer for different laser powers and exposure times to determine the smallest achievable size of the volume pixel (voxel). Figure 6 shows the resulting voxel sizes. For both objectives, the smallest achievable voxel diameters are slightly above 200 nm. The differences in the applied average powers (measured in front of the objectives) between Figs. 6(a) and 6(b) are due to the different aperture sizes of the objectives.

The improvements of the hybrid optics owing to the correction of writing depth dependent aberration become obvious, when applied for volume structuring of the polymer. We fabricated 30 μm high walls on the substrate that act as suspension for lines, written in different heights with constant writing parameters. The resulting line widths in dependence of the writing depth in the polymer can be seen in Fig. 7. The width of lines written with the Zeiss microscope objective [Fig. 7(a)] becomes smaller with increasing polymer thickness. This effect is caused by the spherical aberration that become stronger with increasing writing depth and the consequent drop of the peak intensity in the focal plane. Apart from the changes in structure size, this drop

![Figure 5](image-url)  
**FIG. 5.** Layout of the specifically designed hybrid optics consisting of an aspheric lens (Edmund Optics 47 093), a DOE, and a half ball lens (N-SK11, $r = 2$ mm) working as solid immersion lens.

![Figure 6](image-url)  
**FIG. 6.** Diameter of volume pixel on the substrate surface written with Zeiss Plan-Apochromat (a) and hybrid optics (b). In both cases, the smallest feature sizes are close to 200 nm. The differences in slope and separation between the curves in (a) and (b) are caused by the different aperture sizes of microscope objective and hybrid optics.
of the peak intensity also leads to a smaller degree of polymerization and, thus, to less stable structures (see Sec. II). In contrast, lines with steady widths for constant writing parameters in different writing depths can be fabricated, when the hybrid optics is applied for focusing into the polymer [Fig. 7(b)]. Due to the correction of the refractive index mismatch-induced spherical aberration, the hybrid optics should allow for homogeneous structuring with constant writing parameters over the complete working distance range of 550 μm (experimentally shown for 30 μm).

A further advantage of our optics is the low internal dispersion. The total GVD calculated from the dispersion curves of the optical materials used and the path lengths amounts to 842 fs², which is significantly less than the GVD of microscope objectives with comparable NA in the range of 2000 fs². Figure 8 shows the temporal broadening of the laser pulses in dependence of the initial pulse duration for hybrid optics and microscope objective. This indicates a significant difference for pulse durations shorter than 100 fs.

IV. CONCLUSION

We investigated the influence of the polymer on structure size and presented a focusing concept that build the foundation for three-dimensional two-photon lithography of microstructures with features sizes on the sub-100 nm scale and high aspect-ratios. It could be shown that the newly synthesized ORMOCER® OC-II is suitable for the fabrication of nanostructures with dimension down to 91 nm. We realized a hybrid optics with a high numerical aperture NA = 1.33 for the aberration-free focusing of the laser pulses into the ORMOCER® OC-I. This optics allows for writing of microstructures with heights up to 550 μm without introducing index-mismatched induced aberrations. Owing to the reduced GVD of the hybrid optics, the temporal broadening of the focused laser pulses could be reduced compared to microscope objectives of comparable NA. The hybrid optics can be easily adapted to different polymers by changing the phase function of the DOE and choosing a material with a proper refractive index for the SIL.

The focusing concept in combination with the novel developed ORMOCER®s with their excellent optical properties and biocompatibility enables us to address future micro-optical and biomedical applications.

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